# persistent Müllerian duct syndrome

Persistent Müllerian duct syndrome is a disorder of sexual development that affects males. Males with this disorder have normal male reproductive organs, though they also have a uterus and fallopian tubes, which are female reproductive organs. The uterus and fallopian tubes are derived from a structure called the Müllerian duct during development of the fetus. The Müllerian duct usually breaks down during early development in males, but it is retained in those with persistent Müllerian duct syndrome. Affected individuals have the normal chromosomes of a male (46,XY) and normal external male genitalia.

The first noted signs and symptoms in males with persistent Müllerian duct syndrome are usually undescended testes (cryptorchidism) or soft out-pouchings in the lower abdomen (inguinal hernias). The uterus and fallopian tubes are typically discovered when surgery is performed to treat these conditions.

The testes and female reproductive organs can be located in unusual positions in persistent Müllerian duct syndrome. Occasionally, both testes are undescended (bilateral cryptorchidism) and the uterus is in the pelvis. More often, one testis has descended into the scrotum normally, and one has not. Sometimes, the descended testis pulls the fallopian tube and uterus into the track through which it has descended. This creates a condition called hernia uteri inguinalis, a form of inguinal hernia. In other cases, the undescended testis from the other side of the body is also pulled into the same track, forming an inguinal hernia. This condition, called transverse testicular ectopia, is common in people with persistent Müllerian duct syndrome.

Other effects of persistent Müllerian duct syndrome may include the inability to father children (infertility) or blood in the semen (hematospermia). Also, the undescended testes may break down (degenerate) or develop cancer if left untreated.

# Frequency

Persistent Müllerian duct syndrome is a rare disorder; however, the prevalence of the condition is unknown.

# **Genetic Changes**

Most people with persistent Müllerian duct syndrome have mutations in the *AMH* gene or the *AMHR2* gene. The *AMH* gene provides instructions for making a protein called anti-Müllerian hormone (AMH). The *AMHR2* gene provides instructions for making a protein called AMH receptor type 2.

The AMH protein and AMH receptor type 2 protein are involved in male sex differentiation. All fetuses develop the Müllerian duct, the precursor to female reproductive organs. During development of a male fetus, these two proteins work together to induce breakdown (regression) of the Müllerian duct. Mutations in the *AMH* and *AMHR2* genes lead to nonfunctional proteins that cannot signal for regression of the Müllerian duct. As a result of these mutations, the Müllerian duct persists and goes on to form a uterus and fallopian tubes.

Approximately 45 percent of cases of persistent Müllerian duct syndrome are caused by mutations in the *AMH* gene and are called persistent Müllerian duct syndrome type 1. Approximately 40 percent of cases are caused by mutations in the *AMHR2* gene and are called persistent Müllerian duct syndrome type 2. In the remaining 15 percent of cases, no mutations in the *AMH* and *AMHR2* genes have been identified, and the genes involved in causing the condition are unknown.

#### Inheritance Pattern

This condition is inherited in an autosomal recessive pattern, which means both copies of the gene in each cell have mutations. The parents of an individual with an autosomal recessive condition each carry one copy of the mutated gene, but they typically do not show signs and symptoms of the condition. However, persistent Müllerian duct syndrome affects only males. Females with two mutated copies of the gene do not show signs and symptoms of the condition.

#### Other Names for This Condition

- persistent oviduct syndrome
- PMDS

# **Diagnosis & Management**

## **Genetic Testing**

 Genetic Testing Registry: Persistent Mullerian duct syndrome https://www.ncbi.nlm.nih.gov/gtr/conditions/C1849930/

## Other Diagnosis and Management Resources

 MedlinePlus Encyclopedia: Undescended testicle repair https://medlineplus.gov/ency/article/003002.htm

# General Information from MedlinePlus

- Diagnostic Tests https://medlineplus.gov/diagnostictests.html
- Drug Therapy https://medlineplus.gov/drugtherapy.html

- Genetic Counseling https://medlineplus.gov/geneticcounseling.html
- Palliative Care https://medlineplus.gov/palliativecare.html
- Surgery and Rehabilitation https://medlineplus.gov/surgeryandrehabilitation.html

#### **Additional Information & Resources**

## MedlinePlus

- Encyclopedia: Intersex https://medlineplus.gov/ency/article/001669.htm
- Encyclopedia: Undescended testicle https://medlineplus.gov/ency/article/000973.htm
- Encyclopedia: Undescended testicle repair https://medlineplus.gov/ency/article/003002.htm

#### Genetic and Rare Diseases Information Center

 Persistent Mullerian duct syndrome https://rarediseases.info.nih.gov/diseases/8435/persistent-mullerian-duct-syndrome

#### Additional NIH Resources

 Eunice Kennedy Shriver National Institute of Child Health and Human Development: Infertility https://www.nichd.nih.gov/health/topics/infertility/Pages/default.aspx

#### **Educational Resources**

- Disease InfoSearch: Persistent Mullerian duct syndrome http://www.diseaseinfosearch.org/Persistent+Mullerian+duct+syndrome/5687
- Eunice Kennedy Shriver National Institute of Child Health and Human Development: Infertility https://www.nichd.nih.gov/health/topics/infertility/Pages/default.aspx
- KidsHealth from Nemours: The Facts on Undescended Testicles http://kidshealth.org/en/parents/cryptorchidism.html
- MalaCards: persistent mullerian duct syndrome http://www.malacards.org/card/persistent\_mullerian\_duct\_syndrome

- Orphanet: Persistent Müllerian duct syndrome http://www.orpha.net/consor/cgi-bin/OC\_Exp.php?Lng=EN&Expert=2856
- University of Michigan Health System: Disorders of Sexual Development http://www.med.umich.edu/yourchild/topics/dsd.htm

# Patient Support and Advocacy Resources

- Accord Alliance http://www.accordalliance.org/
- Accord Alliance http://www.accordalliance.org/
- National Organization for Rare Disorders (NORD): Ovotesticular Disorder of Sex Development https://rarediseases.org/rare-diseases/ovotesticular-disorder-of-sex-development/
- The MAGIC Foundation https://www.magicfoundation.org/

# ClinicalTrials.gov

ClinicalTrials.gov https://clinicaltrials.gov/ct2/results?cond=%22persistent+Mullerian+duct+syndrome %22+OR+%2246%2CXY+DSD%22+OR+%22Male+Pseudohermaphroditism %22+OR+%22Male+Pseudohermaphroditisms%22+OR+%2246%2C+XY +Disorders+of+Sex+Development%22+OR+%22Disorders+of+Sex+Development %22

#### Scientific Articles on PubMed

PubMed

https://www.ncbi.nlm.nih.gov/pubmed?term=%28%28persistent+mullerian+duct +syndrome%5BTIAB%5D%29+OR+%28pmds%5BTIAB%5D%29%29+AND+eng lish%5Bla%5D+AND+human%5Bmh%5D+AND+%22last+1440+days%22%5Bdp %5D

#### **OMIM**

 PERSISTENT MULLERIAN DUCT SYNDROME, TYPES I AND II http://omim.org/entry/261550

# **Sources for This Summary**

- Belville C, Van Vlijmen H, Ehrenfels C, Pepinsky B, Rezaie AR, Picard JY, Josso N, di Clemente N, Cate RL. Mutations of the anti-mullerian hormone gene in patients with persistent mullerian duct syndrome: biosynthesis, secretion, and processing of the abnormal proteins and analysis using a three-dimensional model. Mol Endocrinol. 2004 Mar;18(3):708-21. Epub 2003 Dec 12. Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/14673134
- Faure E, Gouédard L, Imbeaud S, Cate R, Picard JY, Josso N, di Clemente N. Mutant isoforms of the anti-Müllerian hormone type II receptor are not expressed at the cell membrane. J Biol Chem. 1996 Nov 29;271(48):30571-5.
  - Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/8940028
- Imbeaud S, Belville C, Messika-Zeitoun L, Rey R, di Clemente N, Josso N, Picard JY. A 27 base-pair deletion of the anti-müllerian type II receptor gene is the most common cause of the persistent müllerian duct syndrome. Hum Mol Genet. 1996 Sep;5(9):1269-77.
  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/8872466
- Josso N, Belville C, di Clemente N, Picard JY. AMH and AMH receptor defects in persistent Müllerian duct syndrome. Hum Reprod Update. 2005 Jul-Aug;11(4):351-6. Epub 2005 May 5. Review.
  - Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/15878900
- Josso N, Picard JY, Imbeaud S, di Clemente N, Rey R. Clinical aspects and molecular genetics of the persistent müllerian duct syndrome. Clin Endocrinol (Oxf). 1997 Aug;47(2):137-44. Review. Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/9302384
- OMIM: PERSISTENT MULLERIAN DUCT SYNDROME, TYPES I AND II http://omim.org/entry/261550
- Rey R. Anti-Müllerian hormone in disorders of sex determination and differentiation. Arq Bras Endocrinol Metabol. 2005 Feb;49(1):26-36. Epub 2006 Mar 16. Review.
   Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/16544032

## Reprinted from Genetics Home Reference:

https://ghr.nlm.nih.gov/condition/persistent-mullerian-duct-syndrome

Reviewed: March 2011 Published: March 21, 2017

Lister Hill National Center for Biomedical Communications U.S. National Library of Medicine National Institutes of Health Department of Health & Human Services